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(54) Title: QUATERNARY AMMONIUM ANTISTATIC COMPOUNDS

(57) Abstract

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Described are certain quaternary ammonium alkyl benzene sulfonate salts and certain quaternary ammonium alkane sulfonate salts, e.g., octyl dimethyl hydroxyethyl ammonium dodecyl benzene sulfonate and octyl dimethyl hydroxyethyl ammonium methane sulfonate which are useful as antistatic agents for synthetic polymer articles.

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OUATERNARY AMMONIUM ANTISTATIC COMPOUNDS

2

DESCRIPTION OF THE INVENTION

The present invention relates to certain novel quaternary ammonium sulfonate compounds and, more particularly, relates to the 6 use of such compounds as internal and external antistatic agents for normally non-conducting organic materials. Still more particularly, 8 the present invention relates to shaped articles of synthetic and naturally ocurring organic polymers that have less than their normal

Organic polymers, e.g., synthetic polymers, are essentially 12 electrical insulators, i.e., non-conductors of electricity. Articles prepared from such polymers tend to develop electrostatic charges upon

14 their surfaces when they are in a dry state and are subjected to friction during their production and finishing or during their

10 tendency to accumulate static charges of electricity.

- 16 handling and use. Such static charges are undesirable for a number of reasons. For example, surface static charges readily attract dust and
- 18 other contaminants, which are unsightly and difficult to clean. Often the contaminants or static charges themselves cause processing or
- 20 handling problems. In certain cases, static charges may accumulate to a level where an unpleasant electric shock is experienced when the
- 22 article is handled. Further, a high level of static charge on a molded part covering sensitive electronic equipment can damage such 24 equipment.

Articles prepared from non-conducting polymers may be surface 26 treated with a finish containing an antistatic agent; however, surface treatments are less desirable than internally incorporated antistatic

- 28 agents due to a wearing away of the applied finish. Conversely, antistatic agents that are incorporated within the shaped article by,
- 30 for example, blending the antistatic agent with the synthetic polymer prior to forming of the article, are susceptible to decomposition as a
- 32 result of the high temperatures used to form the article, e.g., by molding and extrusion. There is, therefore, a continuing need for
- 34 more thermally stable antistatic agents that may be used with synthetic and naturally occurring polymers, and other naturally ccurring materials.

The art has described the use of certain quaternary ammonium 2 p-toluene sulfonate salts as antistatic agents. While some of these salts are relatively thermally stable, e.g., at temperatures of about 4 200°C., their performance as antistatic agents is not as efficacious as desired. In particular, the decay time of a charge applied to an

- 6 article treated with the antistatic agent is too long. It is preferred that such decay times be less than 2 seconds, more
- 8 preferably less than 1 second, and still more preferably less that 0.6 seconds.
- 10. It is an object of the present invention to provide novel antistatic agents for use internally or externally with formed
- 12 synthetic materials, e.g., polymers, and other articles of manufacture to lessen the accumulation thereon of surface static charges.
- It has now been discovered that certain quaternary ammonium sulfonate salts possess both thermal stability and markedly improved
- 16 antistatic properties, i.e., electrostatic buildup and dissipation properties. Such properties may be measured by Federal Test Standard
- 18 101C, Method 4046, which measures the decay time for an applied charge of 5,000 volts. More particularly, the aforesaid quaternary salts may 20 be represented by the following graphic formula I:

$$\begin{bmatrix} R_1 \\ R & N \\ R_2 \end{bmatrix} \bigoplus Y \bigcirc$$
28 (I)

- 30 wherein R is a C_2 - C_{22} alkyl, preferably a C_8 - C_{18} alkyl, and R_1 is selected from the group consisting of C_1 - C_{22} alkyl and an alkyleneoxy
- 32 radical, "Z", that may be represented by the formula, $\{CH_2-C(A)H-0\}_{X}H$, wherein A is hydrogen, methyl or ethyl, and x is an integer of from
- 34 1-5, e.g., hydroxyethyl, hydroxypropyl, hydroxybutyl,
 poly(ethyleneoxy) hydroxyethyl, poly(propyleneoxy) 2-hydroxypropyl,
- 36 and poly(butyleneoxy) 2-hydroxybutyl. Preferably, R_1 is selected from the group consisting of C_1-C_3 alkyl or C_8-C_{18} alkyl, e.g. methyl,

ethyl, n-propyl, isopropyl, octyl, decyl, dodecyl, hexadecyl, and 2 octadecyl, or the alkyleneoxy radical Z wherein A is hydrogen or methyl and x is 1 to 3. Still more preferably, R₁ is a C₁-C₃ alkyl or 4 alkyleneoxy radical Z wherein A is hydrogen and x is 1 to 2. When x is greater than 1, the sulfonate compound may be liquid, which makes 6 it easier to handle.

R₂ in the above graphic formula I is selected from the group 8 consisting of C₁-C₃ alkyl, e.g., methyl, ethyl, n-propyl and isopropyl, and the radical Z, wherein A and x are as defined with 10 respect to R₁. Alternatively R₁ and R₂ may also join together to form a six-membered morpholino group.

- 12 R₃ in the above graphic formula I is a group represented by the alkyleneoxy radical Z, wherein A and x are as defined with respect
- 14 to R_1 ; and Y is the anion, R'SO₃, wherein R' is a C_1-C_{18} alkyl, preferably a C_1-C_2 alkyl, e.g., methyl and ethyl, or a C_8-C_{18}
- 16 alkylphenyl, preferably a $C_{10}-C_{13}$ alkylphenyl. Preferably, the alkylphenyl is a para-alkylphenyl.
- With respect to R, R₁, and R', the term alkyl denotes a univalent, essentially saturated branched or straight chained alkyl
- 20 group. Representative of such alkyl groups include methyl, ethyl, propyl, butyl, pentyl, hexyl, isohexyl, heptyl, octyl, nonyl, decyl,
- 22 undecyl, dodecyl, cocoyl, tridecyl, tetradecyl, pentadecyl, hexadecyl, heptadecyl, octadecyl, soya, eicosyl and the like. When derived from
- 24 naturally occurring materials, the group R, R_1 , and R' may contain a small amount of unsaturation and may be comprised of a mixture of
- 26 alkyl groups. For example, commercially available dodecyl benzene sulfonic acid is a benzene sulfonic acid in which the alkyl
- 28 substituent on the benzene ring is a mixture of $C_{10}-C_{13}$ alkyl groups, the nominal number of carbon atoms being about 12.
- The aforedescribed quaternary ammonium sulfonate salts may be more graphically illustrated by the following graphic formulae II and
- 32 III where R, R_1 , R_2 , and R_3 have the meanings described heretofore with respect to graphic formula I, R_a is a C_1-C_{18} alky1, e.g., C_1-C_2
- 34 alkyl, and R_b is a C_8 - C_{18} alkyl, preferably a C_{10} - C_{13} alkyl. As shown, R_b is a para-alkyl substituent, but it may be an ortho- or 36 meta-substituent.

$$\begin{bmatrix} R_{1} \\ R_{-} \\ R_{-} \\ R_{2} \\ R_{2} \\ R_{3} \end{bmatrix} \xrightarrow{\mathbb{R}_{a}} \begin{bmatrix} R_{1} \\ R_{-} \\ R_{-} \\ R_{2} \\ R_{2} \\ R_{2} \\ R_{3} \end{bmatrix} \xrightarrow{\mathbb{R}_{b}} \mathbb{R}_{b} \xrightarrow{\mathbb{R}_{3}} \mathbb$$

Non-limiting examples of quaternary ammonium sulfonate

10 compounds illustrated by graphic formula I include both the alkane
sulfonate salts, such as methane sulfonate salts, and the alkyl

12 benzene sulfonate salts, such as p-alkyl benzene sulfonate salts, that
are described in Table I. There, R, R₁, R₂, R₃ and R_b are selected

14 from the substituents tabulated therein.

<u>TABLE 1</u> A. <u>Methane Sulfonates</u>

18					
20	Compound/	R	R ₁	R ₂	R ₃
22	1 .	Octy1	Methy1	Methyl	Hydroxyethy1
	2	Octy1	Ethy1	Ethy1	Hydroxyethy1
24	3	Soya	Methy1	Methyl	Hydroxyethy1
	4	Decyl	Decyl	Methy1	Hydroxyethy1
26	5	Octyl	Methyl	Methyl	2-hydroxypropy1
	6	Octy1	Methyl	Methy1	3-hydroxybuty1
28	7	Octy1	4-Morph	olino	Hydroxyethy1
	8	Octy1	Methy1	Hydroxyethy1	Hydroxyethy1
30	9	Decyl	Hydroxyethy1	Hydroxyethyl '	Hydroxyethy1
	10	Octy1	Hydroxyethy1	Hydroxyethy1	Hydroxyethy1
32	11	Octy1	Methyl	Methy1	Poly(ethyleneoxy) hydroxyethyl
34					
	12	Cocoy1	Hydroxyethyl	Hydroxyethy1	Hydroxyethy1
36					

38

16

В.	p-A1kv1	Benzene	<u>Sulfonates</u>

4	Compound/	R	- R ₁	R ₂	R ₃	$R_{\mathbf{b}}$
6					. —	
	1	Octy1	Methy1	Methyl	Hydroxyethy1	Dodecy1
8	2	0cty1	Methy1	Methyl	2-hydroxypropyl	Dodecy1
	3	0ctyl	Methy1	Methyl	3-hydroxybutyl	Dodecyl
10	4	Soya	Methy1	Methy1	Hydroxyethy1	Dodecyl
	5	0ctyl	0cty1	Methy1	Hydroxyethy1	Dodecy1
12	. 6	Decy1	Decy1	Methyl	Hydroxyethyl	Dodecy1
14	7	Octy1	Methy1	Methy1	Poly(ethyleneoxy) hydroxyethyl	Dodecy1

16 Compound 1 in Table 1, Part A may be named octyl dimethyl 2-hydroxyethyl ammonium methane sulfonate (where Y is methane

- 18 sulfonate). Compound 1 in Table 1, Part B may be named octyl dimethyl 2-hydroxyethyl ammonium p-dodecyl benzene sulfonate (where Y is
- 20 dodecyl benzene sulfonate). Other sulfonate salts of the compounds from Table 1 may be similarily named by utilizing the appropriate
- 22 IUPAC radical name for the designated substituents.

The quaternary ammonium sulfonate salts of graphic formula I

- 24 may be used to minimize the accumulation of static electricity on non-conducting articles, e.g., articles prepared from synthetic
- 26 polymers, by applying to the surface of or incorporating within the article effective antistatic amounts of at least one of the salts.
- 28 Generally, the compounds of graphic formula I may be incorporated within the article in amounts of between about 0.5 and about 20 weight
- 30 percent and preferably are used in amounts of between about 2 and about 6 weight percent, based on the weight of the dry, untreated
- 32 article. When associated with a finish composition, the compounds of graphic formula I are generally present in amounts of from about 0.1
- 34 to about 2, e.g., 0.5 to 1, weight percent.

When compounded into a shaped article of a synthetic polymer, 36 the antistatic quaternary ammonium sulfonate salt will migrate or "blo m" to the surface of th article to provide an antistatic coating thereon. Such a coating is more permanent than an ext rnally
2 (topically) applied coating since the latter can be removed by wear,
wiping, washing, handling, movement in transit, etc. In contrast, if
4 the migrated layer of an internally compounded antistatic agent should
be removed during handling or processing, a new antistatic layer will
6 bloom to the surface.

To apply the antistatic compound topically to the surface of 8 an article, the antistatic compound is dissolved or dispersed in water, lower alkanol, e.g., a C₁-C₄ alcohol, lubricating oil, 10 polymeric coating, other organic solvent, etc. and the resulting "finish" containing the desired amount of antistatic compound applied 12 to the surface using conventional coating techniques, e.g., spraying, dipping, wiping, etc., thereby to deposit an effective amount of 14 antistatic agent on the surface of the article.

In general, the antistatic compounds of graphic formula I are thermally stable at temperatures in excess of 200°C. When heated to temperatures up to about 275°C., weight losses of less than 5 weight percent have been observed for compounds such as octyl dimethyl hydroxyethyl ammonium methane sulfonate. The compounds may be solids, 20 liquids or low melting waxes, depending on the number of carbons in the R, R₁, R_a and R_b alkyl groups. At the same time, these compounds are more antistatic, i.e., exhibit shorter decay times, than sulfonate salts currently in commercial use, e.g., p-toluene sulfonate salts.

When antistatic compounds are incorporated internally into an article, the compound may be mixed in antistatic amounts with the synthetic polymer or other material by conventional blending or mixing equipment, e.g., Banbury mixers or other rubber and plastic processing equipment, and the mixture formed into the article, e.g., by extrusion

or other molding procedure. Alternatively, a master batch of the
30 polymer and antistatic compound may be prepared and the master batch
added in antistatic amounts to synthetic polymer that is to be formed
32 into an article, thereby to provide the desired antistatic amount of
antistatic compound within the article. A master batch may contain

34 between about 10 and about 25 percent by weight of the antistatic compound.

The compounds of graphic formula I may be used with

- 2 conventional synthetic polymers utilized to prepare formed articles.

 The compatibility of the antistatic compounds with a particular
- 4 synthetic polymer may be readily determined by those skilled in the art. Antistatic compounds of graphic formula I may be used in a wide
- 6 spectrum of substrate shapes, such as fibers (woven and nonwoven), sheets, films and molded or extruded articles. Such articles may be
- 8 prepared from thermoplastic or thermosetting polymers or copolymers (including terpolymers).
- Non-limiting examples of synthetic polymers from which formed articles may be prepared include polyolefins, such as polyethylene,
- 12 polypropylene and polyisobutylene, styrene resins such as polystyrene, poly(chlorostyrene), styrene-acrylonitrile copolymers, poly(styrene-
- 14 acrylonitrile-butadiene) terpolymers (ABS resins) and high impact polystyrene (HIPS), polyesters such as poly(methylacrylate),
- 16 poly(methylmethacrylate) and poly(vinylacetate), ethylene glycolterephthalic acid polymers, polycarbonates, polyamides such as nylon
- 18 and Kevlar® type polyamides, polyacetals such as poly(vinylbutyral), phenol-formaldehyde resins, vinyl resins, such as poly(vinyl
- 20 chloride), poly(vinylidene chloride), polytrifluorochloroethylene, copolymers of vinyl chloride with vinyl acetate, vinylidene chloride,
- 22 or acrylonitrile, polyurethanes, and poly(phenylene ether) resins.

 Mixtures of the aforesaid polymers may also be used, e.g., polymer

 24 alloys.
- In addition, the compounds of graphic formula I may be used
- 26 with natural materials or mixtures of natural and synthetic materials, e.g., rayon, acetate, rayon-cellulosic materials such as cellulose
- 28 acetate-propionate, cellulose-butyrate, cotton, linen, jute, ramie, wool, mohair and glass, e.g., fiberglass and fiberglass insulation.
- 30 The textile materials may take any form, including individual fibers, woven material such as fabrics, cloth, carpets, rugs and upholstery
- 32 and non-woven materials such as felts, bats and mats. In the case of fiberglass strand or fiberglass insulation, the compounds of graphic
- 34 formula I may be applied topically as a finish or as part of a sizing composition.

The compounds of graphic formula I may be prepared by
2 reacting approximately equal molar amounts of the tertiary amine and
an ester of alkane sulfonic acid or alkyl benzene sulfonic acid in the

- 4 presence of a solvent, e.g., water, lower alkanols, e.g., C₁-C₄ alkanols, or acetone, at tempatures of from about 25°C. to about
- 6 250°C., e.g., from about 65 to 80°C., under a slight positive nitrogen pressure. Preferably, the compounds of graphic formula I may be
- 8 prepared more readily by reacting the corresponding tertiary amine, corresponding sulfonic acid, e.g., methane sulfonic acid, and alkylene
- 10 oxide, e.g., ethylene oxide, in accordance with the following equation IV.

12

14
$$R - N + R'SO_{3}H + CH_{2}-CH_{2}-O \rightarrow R-N-CH_{2}CH_{2}OH R'SO_{3}C$$
16
$$R_{2} R_{2} R_{2} (IV)$$

Quaternization of tertiary amines with alkylene oxide is
20 preferably performed in the presence of a small amount of amine
catalyst and at elevated pressures, the later accelerating the take-up
22 of the alkylene oxide. Quaternary amines containing poly(alkylene
oxide), e.g., poly(ethylene oxide) groups may preferably be prepared
24 in the presence of a slight excess of the amine, an excess of alkylene
oxide and at elevated pressures, which may be from about 10 to about
26 100 pounds per square inch (.07-0.7 MPa).

The present invention is more particularly described in the 28 following examples which are intended as illustrative only since numerous modifications and variations will be apparent to those 30 skilled in the art.

32 Example 1

118.0 grams (0.75 moles) of n-octyl dimethylamine were

34 neutralized with a solution of 72.0 grams (0.75 moles) of methane
sulfonic acid in 100 milliliters of water. The temperature of the

resulting mixture rose to about 80-90°C. and thereafter was cooled to 2 about 40°C. The reaction mixture was charged to an autoclave and, after thoroughly flushing the system with nitrogen, the autoclave was 4 sealed and pressurized with nitrogen to 20 pounds per square inch (psig) (0.1 MPa). The contents of the autoclave were heated to about 6 70°C. and 62.0 grams (1.4 moles) of ethylene oxide charged to the autoclave. The contents of the autoclave were stirred for 1 hour 8 while maintaining the temperature at 70°C. Thereafter, the autoclave was flushed with nitrogen and the reaction mixture cooled. The pH of 10 the cooled solution in the autoclave was determined to be about 10.5 and subsequently adjusted to about 4.7 using methane sulfonic acid and 12 a 50 percent sodium hydroxide aqueous solution. 50.0 grams of the solution in the autoclave were stripped under vacuum; the residue 14 dissolved in an acetone-ether solution; and the solution decolorized with charcoal. The decolorized mixture was filtered and concentrated 16 to obtain 31.5 grams of a clear liquid that solidified on standing. The product, n-octyl dimethyl hydroxyethyl ammonium methane sulfonate, 18 was confirmed by discharge ionization secondary ion mass spectrometry

20

(DISIMS).

Example 2

62.0 grams (0.28 moles) of n-octyl diethanol amine were 22 neutralized with a solution of 27.0 grams (0.28 moles) of methane 24 sulfonic acid in 100 milliliters of water. The pH of the mixture was determined to be about 5.3 and was adjusted subsequently to 6.3 with 26 octyl dimethyl amine. 50 milliliters of water were added to the neutralized mixture and the resulting solution charged to an 28 autoclave, which was first flushed with nitrogen and then pressurized with nitrogen to 20 psig (0.1 MPa). The contents of the autoclave 30 were heated to 65°C. and 25.0 grams (0.37 moles) of ethylene oxide added thereto over a period of 30 minutes. After addition of the 32 ethylene oxide, the nitrogen pressure was increased to 100 pounds psig (0.7 MFa) and the reaction mixture stirred for 2 hours. The reaction 34 mixture was then cooled, the autoclave flushed with nitrogen, and the contents of the autoclave stripped under vacuum to obtain 108 grams of 36 a viscous liquid. The product, n-octyl tris(hydroxyethyl) ammonium methane sulfonate, was confirmed by DISIMS spectrometry.

2 Example 3

dodecyl benzene sulfonate.

To a solution of 123.3 grams (0.41 moles) of soya dimethyl

4 amine in 200 milliliters of isopropanol were added with stirring 132.0 grams of dodecyl benzene sulfonic acid. The temperature of the

6 mixture rose to about 50°C. The pH of the mixture was determined to be about 1.8. The mixture was charged to an autoclave, which was

8 sealed and pressurized with nitrogren to 20 psig (0.1 MPa). The mixture was heated to 70°C. and 35.0 grams (0.80 moles) of ethylene

10 oxide added to the reactor over a period of 30 minutes. The mixture was stirred overnight while maintaining it at a temperature of 70°C.

12 The pressure was then released and the contents of the autoclave cooled. Isopropanol was stripped from the crude reaction product on a

14 rotary evaporator to obtain 252 grams of a slight yellow paste. The structure of the product was determined by mass spectroscopy to be a

16 mixture corresponding to n-soya dimethyl hydroxyethyl ammonium

18

Example 4

To a solution of 160.0 grams (0.5 moles) of dodecyl benzene sulfonic acid in 150 milliliters of ethanol were added with stirring 22 78.3 grams (0.5 moles) of octyl dimethyl amine. The mixture was charged to an autoclave which was pressurized with nitrogen to a 24 pressure of 20 psig (0.1 MPa.) The mixture was heated to 65°C. and 28.0 grams (0.64 moles) of ethylene oxide added with stirring to the 26 autoclave over a period of 30 minutes. The mixture was then stirred for 2 hours while maintaining it at a temperature of 65°C. The 28 pressure in the autoclave was released, the contents cooled and the ethanol solvent stripped in vacuo to obtain 244.0 grams of a thick 30 liquid.

Since no weight gain was observed, the liquid product was
32 redissolved in 200 milliliters of isopropanol and charged to an
autoclave. The contents of the autoclave were warmed to 80°C. and the
34 autoclave pressurized with nitrogen to 20 psig (0.1 MPa). Ethylene
oxide (44.0 grams, 1.0 mole) was added to the autoclave over a period
36 of 30 minutes and the autoclave contents stirred overnight while

maintaining the contents of the autoclave at a temperature of 80°C.

2 and at a pressure of 100 psig (0.7 MPa). The pressure was r leased and the reaction product cooled. The isopropanol solvent was stripped 4 in vacuo to obtain 262.0 grams of a light yellow gel. This product was found to be octyl dimethyl hydroxyethyl ammonium dodecyl benzene 6 sulfonate by mass spectroscopy.

Example 5

To a solution of 160.0 grams of dodecyl benzene sulfonic acid 10 in 100 milliliters of isopropanol were added 79.5 grams of octyl dimethyl amine. The mixture was charged to an autoclave which was 12 pressurized with nitrogen to a pressure of 20 psig (0.1 MPa). The mixture was warmed to 80°C. and 35.0 grams of propylene oxide added 14 with stirring to the autoclave over a period of about 30 minutes. The mixture was stirred overnight while maintaining it at a temperature of 16 80°C. In the morning the autoclave pressure had dropped to 10 psig (0.07 MPa), which indicated that the autoclave seal had a leak. 18 The solvent was stripped from the mixture and the residue (256.8 grams) redissolved in 150 milliliters of isopropanol and the 20 solution charged to the autoclave. Propylene oxide (30.0 grams) was added to the autoclave, the contents warmed to 80°C. and the autoclave 22 pressurized to 30 psig (0.2 MPa) with nitrogen. The pressure dropped to 20 psig (0.1 MPa) in 1 hour. The mixture was stirred overnight at 24 a temperature of 60°C. In the morning the pressure was found to be about 5 psig (0.03 MPa). The mixture was cooled and the solvent 26 stripped on a rotary evaporator. The residue (270.5 grams) was the product octyl dimethyl 2-hydroxypropyl ammonium dodecyl benzene

Example 6 (Comparative)

28 sulfonate, which was confirmed by mass spectroscopy.

157.3 grams of octyl dimethyl amine were charged to an
32 autoclave and neutralized with a solution of 189.5 grams of p-toluene
sulfonic acid in 100 milliliters of water. The temperature of the
34 mixture was about 60°C. at the end of the neutralization step. The
autoclave was sealed, flushed twice with nitrogen and pressurized to

25 psig (0.2 MPa) with nitrogen. The mixture was warmed to 75-80°C. 2 and 46.0 grams of ethylene oxide added to the autoclave over about 1 hour. The mixture was stirred at 70°C. for 1 hour.

The pressure in the autoclave was released and the product recovered. A portion of the product was stripped in vacuo and the 6 residue washed with ether. A white powder product was obtained by filtration. Mass spectroscopy confirmed the product to be octyl 8 dimethyl hydroxyethyl ammonium p-toluene sulfonate.

10 . Example 7

An aqueous solution containing 0.09 weight percent of the 12 compound of Example 1 was prepared. A 3 1/2 inch (8.9 cm) x 5 inch (12.7 cm) swatch of an acrylic fabric obtained from Testfabrics Inc.

- 14 was immersed in 100 grams of the aqueous solution for 30 seconds.

 Excess solution was squeezed from the swatch using an Atlas laboratory
- 16 wringer. The swatches were air dried and then conditioned for 24 hours in a controlled humidity chamber maintained at 15 percent
- 18 relative humidity. This procedure was repeated with the compounds of Examples 2, 4, 5 and 6, and with the commercial antistatic compound
- 20 Hexcel®106G. Electrostatic properties of the swatches were measured in accordance with Federal Test Standard 101C, Nethod 4046 by applying
- 22 a charge of 5,000 volts to each of the acrylic fabric swatches and measuring the time in seconds required for the charge to decay to 0
- 24 volts with an Electrotech Systems static decay meter, Model 406C. Results are tabulated in Table 2.

26

Table 2

28

Example (Compound) 1 2 4 5 6 Hexcel 106G*

30

Decay Time (sec) 0.3 0.5 0.6 0.9 10 1.7

32 * octyl methyl di(hydroxyethyl) ammonium p-toluene sulfonate

The data of table 2 show that compounds of graphic formula I, e.g., compounds of Examples 1,2,4 and 5 have excellent antistatic

36 properties - exhibiting superior decay times compared to the comp und

of Example 6 (octyl dimethyl hydroxyethyl p-toluene sulfonate) and 2 significantly improved decay times compared to the commercial antistatic product Hexcel® 106G.

4

Example 8

Noryl® N-190 modified poly(phenylene oxide) resin and 2
weight percent of antistatic compound corresponding to Example 1 were
8 blended for 5 minutes in a Brabender mixer (100 rpm) at 210°C. The
resulting mixture was compression molded at 440°F. (227°C.) into a
10 flat plaque about 55 mils thick. The aforesaid procedure was repeated
except that 5 weight precent of the antistatic compound was used.

Five inch (12.7 cm) square discs cut from each of the plaques were conditioned for 24 hours in a controlled humidity chamber

14 maintained at 15 percent relative humidity. The surface resistivity

and electrostatic property (decay time) of the discs were measured 16 using the test method described in Example 7. Results are tabulated in Table 3.

18

Example 9

The procedure of Example 8 was repeated using the compound of Example 4. Results are tabulated in Table 4.

22

Example 10

High impact polystyrene (HIPS-Mobil polystyrene 4226) and 2
weight percent of antistatic compound corresponding to Example 1 were
blended for 3 minutes in a Brabender mixer (100 rpm) at 210°C. The
resulting mixture was compression molded at 420°F. (216°C.) into a

28 flat plaque about 55 mils thick. The procedure was repeated except
that 5 weight percent of the antistatic compound was used. The

30 surface resistivity and electrostatic property of each of the plaques
were tested in the manner described in Example 8. Results are

34

32 tabulated in Table 3.

Example 11

The procedure of Example 10 was repeated using the compound 36 of Example 4. Results are tabulated in Table 4.

Example 12

A poly(vinyl chloride) plastisol resin was prepared in the 4 following menner. 92 parts of Geon 173 dispersion resin and 8 parts of Borden 260 ss blending resin were mixed in a blender. 49 parts of

- 6 dioctyl phthalate and 4 parts of Synpron 1363 thermal stabilizer were mixed and the mixture added with stirring slowly to the blender.
- 8 After all of the dioctyl phthalate/stabilizer blend had been added, the resulting mixture was stirred for an additional 5 minutes. 0.75
- 10 grams of the antistatic compound corresponding to Example 1 was added to 14.25 grams of the plastisol resin and the mixture stirred well. A
- 12 10 mil film was drawn down on a 7 inch (17.8 cm) square piece of cardboard and the film cured at 193°C. for 2 minutes. The surface
- 14 resistivity and electrostatic property of the film were measured using the method described in Example 7. Results are tabulated in Table 3.

16

Example 13

The procedure of Example 12 was repeated using the compound of Example 4. Results are tabulated in Table 4.

20

22

TABLE 3 PLASTIC RESIN

24	NORYL		HIPS		PVC _PLASTISOL
Antistat, Wt% 26	_2_	<u> 5 </u>	_2_	_5_	_5_
Decay Time, Secs	1.2	0.2	0.9	0.03	0.07
28 Resistivity, ohm/sq	3×10^{10}	5x10 ¹¹	2×10^{11}	3x10 ⁹	1.5x10 ¹⁰

The data of Table 3 shows that the compound of Example 1 exhibits excellent antistatic properties — showing decay times of less than 1 second for each of the plastic resins at the 5 percent level.

TABLE 4

2	PLASTIC	RESIN

L	NORYL		HIPS		PVC PLASTISOL	
Antistat, Wt%	2	_5_	_2_	_5_	_5_	
Decay Time, secs	1.8	0.7		1.1	0.5	
8 Resistivity, ohm/sq	8×10^{11}	$2x10^{11}$	1×10^{12}	$4x10^{11}$	5x10 ¹⁰	

The data of Table 4 show that the compound of Example 4 has good antistatic properties at a 2 weight percent level and very good 12 antistatic properties at a 5 weight percent level - showing decay times of about 1 second or less for the later level of compounding.

Example 14

To a solution of 48.0 grams of 98% methane sulfonic acid in 150 milliliters of water were added with stirring 79.0 grams of octyl 18 dimethyl amine. The temperature of the reaction was maintained at 60-65°C. The pH of the resulting solution was determined to be 1.5 20 and was subsequently adjusted to 6 with additional octyl dimethyl amine. The solution was charged to an autoclave, which was 22 pressurized with nitrogen to 30 psig (0.2 MPa). The contents in the autoclave were heated to 70°C. and 48.0 grams of ethylene oxide were 24 added over a period of 30 minutes. Thereafter the reaction mixture was stirred at 70°C. for 1.5 hours. The pressure in the autoclave was 26 released and the reaction product stripped in vacuo to obtain 174.0 grams of a liquid product which remained liquid even on prolonged 28 storage. The product was identified by mass spectroscopy to correspond to the products C8H17N (CH3)2 (CH2CH20)xH CH3SO3, wherein x

32 Example 15

30 was 1,2 and 3.

A solution of 68.5 grams of 70% methane sulfonic acid in 150
34 milliliters of water were neutralized with 107.0 grams of dodecyl
dimethyl amine at 80-90°C. The pH of the neutralized solution was
36 determined to be 1.8, which was subsequently adjusted to 5 with
additional dodecyl dimethyl amine. The resulting solution was charged

- to an autoclave, which was pressurized with nitrogen t 20 psig (0.1
- 2 MPa), and the solution heated to 70°C. 68.0 grams of ethylene oxide was added to the autoclave over 45 minutes. After addition of the
- 4 ethylene oxide, the reactor contents were stirred for one hour while maintaining it at 70°C. Pressure in the autoclave was released and
- 6 the liquid product stripped in vacuo to obtain 224.0 grams of a liquid product. Mass spectroscopy showed the product to be a mixture of the
- 8 mono- and diethoxylated quaternary amine, i.e., $C_{12}H_{25}N(CH_3)_2$ ($CH_2CH_2O)_xH$, wherein x is 1 and 2.
- Although the present invention has been described with reference to specific details of certain embodiments thereof, it is
- 12 not intended that such details should be regarded as limitations upon the scope of the invention except as and to the extent that they are
- 14 included in the accompanying claims.

CLAIMS

2

1. A quaternary ammonium compound represented by the graphic 4 formula:

$$\begin{bmatrix} R_1 \\ R & \\ R_2 \end{bmatrix} \oplus R_b - So_3 \oplus$$

wherein R is a C₂-C₂₂ alkyl, R₁ is selected from the group consisting
12 of C₁-C₂₂ alkyl and an alkyleneoxy radical represented by the formula

†CH₂-C(A)H-O]_XH, R₂ is selected from the group consisting of C₁-C₃
14 alkyl, and the alkyleneoxy radical, †CH₂-C(A)H-O]_XH, or R₂ may join

together with R₁ to form a morpholino group, R₃ is an alkyleneoxy
16 radical represented by the formula †CH₂-C(A)H-O]_XH, R_b is a C₈-C₁₈

alkyl, A is selected from the group consisting of hydrogen, methyl and
18 ethyl, and x is an integer of from 1 to 5.

- 20 2. The quaternary ammonium compound of claim 1 wherein R is a C₈-C₁₈ alkyl, R₁ is selected from the group consisting of C₁-C₃
 22 alkyl, C₈-C₁₈ alkyl and the alkyleneoxy radical, fCH₂-C(A)H-O]_xH, R₂ is selected from the group consisting of C₁-C₃ alkyl and
 24 fCH₂-C(A)H-O]_xH, R₃ is fCH₂-C(A)H-O]_xH, R_b is a C₁₀-C₁₃ alkyl, A is hydrogen or methyl and x is an integer of from 1 to 3.
- 3. The quaternary ammonium compound of claim 2 wherein R_1 is 28 selected from the group consisting of C_1-C_3 alkyl and ${TCH_2-C(A)H-O]_xH}$.

30

4. The quaternary ammonium compound of claim 2 wherein R₁
32 and R₂ are each selected from the group consisting of C₁-C₂ alkyl and {CH₂-C(A)H-O]_xH, R₃ is {CH₂-C(A)H-O]_xH, and A is hydrogen.

•

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5. The quaternary ammonium compound of claim 2 wherein R_1 2 and R_2 are each C_1 - C_2 alkyl and R_3 is the radical, $\{CH_2-C(A)H-O\}_xH$, and A is hydrogen or methyl.

4

- 6. The quaternary ammonium compound of claim 2 wherein R_1 , 6 R_2 and R_3 are each $\{CH_2-C(A)H-O\}_xH$, and A is hydrogen.
- 7. The quaternary ammonium compound of claim 2 wherein R_1 is C_1-C_2 alkyl and R_2 and R_3 are each $\{CH_2-C(A)H-O\}_xH$, and A is hydrogen.
- 8. The quaternary ammonium compound of claim 5 wherein R is 12 octy1, R_1 and R_2 are each methyl, R_3 is the radical, $-(CH_2-C(A)H-O]_xH$, A is hydrogen or methyl, x is an integer of from 1 to 3, and R_b is a 14 $C_{10}-C_{13}$ alkyl.
- 16 9. The quaternary ammonium compound of claim 8 wherein A is hydrogen and $R_{\rm b}$ is dodecyl.

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- 10. The quaternary ammonium compound of claim 8 wherein A is 20 methyl and $R_{\rm h}$ is dodecyl.
- 22 11. The quaternary ammonium compound of claim 5 wherein R is soya, R₁ and R₂ are each methyl, R₃ is the radical, †CH₂-C(A)H-O]_XH, A 24 is hydrogen, x is an integer of from 1 to 3, and R_b is dodecyl.
- 12. The quaternary ammonium compound of claim 5 wherein R is octyl, R₁ is methyl, R₂ and R₃ are each the radical, [CH₂-C(A)H-O]_xH, 28 wherein A is hydrogen, x is an integer of from 1 to 3 and R_b is a C₁₀-C₁₃ alkyl.

30

- 13. The quaternary ammonium compound of claim 12 wherein $R_{\mbox{\scriptsize b}}$ 32 is dodecyl.
- 34 14. A quaternary ammonium compound represented by the graphic formula:

20

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$$\begin{bmatrix} R_1 \\ R & \\ R_2 \end{bmatrix} \oplus R_a so_3 \in \mathbb{R}$$

8 wherein R is octyl, R₁ and R₂ are each selected from the group consisting of C₁-C₂ alkyl and the radical, fCH₂-C(A)H-O]_XH, R₃ is the 10 radical, fCH₂-C(A)H-O]_XH, R_a is C₁-C₂ alkyl, A is selected from the group consisting of hydrogen, methyl and ethyl, and x is an integer of 12 from 1 to 3.

14 15. The quaternary ammonium compound of claim 14 wherein R_1 and R_2 are each methyl, R_a is methyl, and A is hydrogen.

16. The quaternary ammonium compound of claim 14 wherein R_1 18 is methyl, R_2 and R_3 are each the radical, $\{CH_2-C(A)H-O\}_xH$, R_a is methyl and A is hydrogen.

17. The quaternary ammonium compound of claim 14 wherein R_1 , 22 R_2 and R_3 are each the radical, $\{CH_2-C(A)H-0\}_xH$, R_a is methyl and A is hydrogen.

18. An organic article having antistatic properties

26 comprising an organic material which in a dry state has a tendency to accumulate static charges of electricity thereon and an antistatically 28 effective amount of a quaternary ammonium compound represented by the graphic formula:

30 $\begin{bmatrix}
R_1 \\
R_1 \\
R_2
\end{bmatrix} \oplus R' \text{ so } 3^{\Theta}$ 36

wherein R is a C_2 - C_{22} alkyl, R_1 is selected from the group consisting

- of C_1-C_{22} alkyl and an alkyleneoxy radical represented by the formula 2 $\{CH_2-C(A)H-0\}_xH$, R_2 is selected from the group consisting of C_1-C_3 alkyl and the alkyleneoxy radical, $\{CH_2-C(A)H-0\}_xH$, or R_2 may join
- 4 together with R_1 to form a morpholino group, R_3 is an alkyleneoxy radical represented by the formula, $\{CH_2-C(A)H-0\}_xH$, R' is selected
- 6 from the group consisting of C_1 - C_{18} alkyl and C_8 - C_{18} alkylphenyl, A is selected from the group consisting of hydrogen, methyl and ethyl, and 8 x is an integer of from 1 to 5.
- 19. The article of claim 18 wherein R is a C₈-C₁₈ alkyl, R₁ is a C₁-C₃ alkyl, C₈-C₁₈ alkyl, or the radical, -{CH₂-C(A)H-O]_xH, R₂ is 12 selected from the group consisting of C₁-C₃ alkyl and the radical, -{CH₂-C(A)H-O]_xH, A is hydrogen or methyl and x is an integer of from 1 to 3.
- 16 20. The article of claim 19 wherein the article is a synthetic polymer.

- 21. The article of claim 20 wherein the synthetic polymer is 20 selected from the group consisting of polyolefins, styrene resins, polyesters, ethylene gylcol-terephthalic acid polymers,
- 22 polycarbonates, polyamides, polyacetals, vinyl resins, poly(phenylene ether) resins, and mixtures of such polymers.

24

22. The article of claim 21 wherein the quaternary ammonium 26 compound is associated with the article in amounts of between about 0.5 and about 20 weight percent.

28

- 23. The organic article of claim 21 wherein R is octyl, R₁
 30 and R₂ are each C₁-C₃ alkyl or the radical, +CH₂-C(A)H-O]_xH, R₃ is the radical +CH₂-C(A)H-O]_xH, R' is C₁-C₂ alkyl or C₁₀-C₁₃ alkylphenyl, A
 32 is hydrogen or methyl and x is an integer of from 1 to 3.
- 34 24. The organic article of claim 23 wherein R_1 and R_2 are each methyl and n is 1.

36

25. A method for treating an article formed of a material 2 which in a dry state has a tendency to accumulate static charges of electricity thereon, which comprises applying to the surface of said 4 article a finish composition containing from about 0.1 to about 2 weight percent of a quaternary ammonium compound represented by the 6 graphic formula:

$$\begin{bmatrix} R_1 \\ R_2 \end{bmatrix} \oplus R'so_3 \ominus$$

wherein R is a C₂-C₂₂ alkyl, R₁ is selected from the group consisting 14 of C₁-C₂₂ alkyl and an alkyleneoxy radical represented by the formula $\{CH_2-C(A)H-O\}_xH$, R₂ is selected from the group consisting of C₁-C₃ 16 alkyl and the radical, $\{CH_2-C(A)H-O\}_xH$, or R₂ may join together with R₁ to form a morpholino group, R₃ is an alkyleneoxy radical 18 represented by the formula, $\{CH_2-C(A)H-O\}_xH$, R' is selected from the group consisting of C₁-C₁₈ alkyl and C₈-C₁₈ alkyl phenyl, A is 20 selected from the group consisting of hydrogen, methyl and ethyl, and x is an integer of from 1 to 5.

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26. The method of claim 25 wherein R is a C₈-C₁₈ alkyl, R₁ is 24 a C₁-C₃ alkyl, C₈-C₁₈ alkyl, or the radical, {CH₂-C(A)H-O]_xH, R₂ is selected from the group consisting of C₁-C₃ alkyl and the radical, 26 {CH₂-C(A)H-O]_xH, A is hydrogen or methyl and x is an integer of from 1 to 3.

28

- 27. The method of claim 26 wherein the article is a synthetic 30 polymer selected from the group consisting of polyolefins, styrene resins, polyesters, ethylene gylcol-terephthalic acid polymers, 32 polycarbonates, polyamides, polyacetals, vinyl resins and poly(phenylene ether) resins, and mixtures of such polymers. 34
- 28. A synthetic polymer composition having antistatic
 36 properties conprising a synthetic polymer having incorporated therein

an antistatically effective amount of a quaternary ammonium compound 2 represented by the graphic formula:

$$\begin{bmatrix} R_1 \\ R & R_3 \end{bmatrix} \oplus R \cdot So_3 \ominus$$

- 10 wherein R is a C_2 - C_{22} alkyl, R_1 is selected from the group consisting of C_1 - C_{22} alkyl and an alkyleneoxy radical represented by the formula
- 12 $_{\rm c}$ = C(A)H-0] $_{\rm x}$ H, R $_{\rm 2}$ is selected from the group consisting of C $_{\rm 1}$ -C $_{\rm 3}$ alkyl and the radical, $_{\rm 1}$ CH $_{\rm 2}$ -C(A)H-0] $_{\rm x}$ H, or R $_{\rm 2}$ may join together with
- 14 R_1 to form a morpholino group, R_3 is a radical represented by the formula, $-\text{CH}_2-\text{C(A)H-Ol}_x\text{H}$, R' is selected from the group consisting of
- 16 C₁-C₁₈ alkyl and C₈-C₁₈ alkyl phenyl, n is a number of from 1 to 3 and x is a number of from 1 to 5.

18

- 29. The polymer composition of claim 28 wherein the synthetic
- 20 polymer is selected from the group consisting of polyolefins, styrene resins, polyesters, ethylene gylcol-terephthalic acid polymers,
- 22 polycarbonates, polyamides, polyacetals, vinyl resins, poly(phenylene ether) resins, and mixtures of such polymers.

24

30. The polymer composition of claim 29 wherein the 26 quaternary ammonium compound is incorporated into the synthetic polymer in amounts of between about 0.5 and about 20 weight percent.

28

- 31. The polymer composition of claim 30 wherein R is octyl,
- 30 R₁ and R₂ are each C₁-C₃ alkyl or the radical, $\{CH_2-C(A)H-O\}_xH$, R₃ is the radical, $\{CH_2-C(A)H-O\}_xH$, R' is C₁-C₂ alkyl or C₁₀-C₁₃
- 32 alkylphenyl, A is hydrogen or methyl and x is an integer of from 1 to 3.

34

32. The polymer composition of claim 31 wherein $\rm R_{\mbox{\scriptsize 1}}$ and $\rm R_{\mbox{\scriptsize 2}}$ are 36 each methyl and A is hydrogen.

2 33. A quaternary ammonium compound represented by the graphic formula:

 $\begin{bmatrix}
R_1 \\
R \\
R \\
R_2
\end{bmatrix} \oplus Raso_3 = Ras$

wherein R is selected from the group consisting of soya, cocoyl and 12 decyl, R₁ and R₂ are each selected from the group consisting of C₁-C₂ alkyl and the radical, {CH₂-C(A)H-O]_xH, R₃ is the radical 14 {CH₂-C(A)H-O]_xH, Ra is C₁-C₂ alkyl, A is selected from the group consisting of hydrogen, methyl and ethyl, and x is an integer of from 16 1 to 3.

18 34. The quaternary ammonium compound of claim 33 wherein R is soya, R_1 and R_2 are each methyl, Ra is methyl and A is hydrogen.